PM2.5 Cation Analysis Revision 5 Date: August 14, 2003 Page 1 of 10

Standard Operating Procedure for PM2.5 Cation Analysis

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TABLE OF CONTENTS

Section	<u>n</u>		<u>Page</u>
1.0	Proced 1.1 1.2 1.3 1.4 1.5 1.6 1.7 1.8 1.9 1.10 1.11 1.12	Purpose and Applicability Summary of Method Health and Safety Warnings Cautions Interferences Personnel Qualifications Apparatus and Materials Reagents Calibration Sample Collection Sample Handling Filter Extraction Procedure 1.12.1 Nylon Filters 1.12.2 Teflon Filters IC Procedure Calculations and Data Reduction	344556777
2.0		Control and Quality Assurance	
3.0	Referen	nce	10
		LIST OF TABLES	
<u>Table</u>]	Page
1. Acc	uracy V	Values for QA/QC Samples and Spiked PM2.5 Filter Extracts	3
2. Prec	cision V	alues for PM _{2.5} Filter Extracts, Extracting Solution, and QC Samples	4
3. Prep	paration	of Ammonium Ion Calibration Standards	6

Page 3 of 10

Standard Operating Procedure for PM2.5 Cation Analysis

1.0 Procedural Section

1.1 Purpose and Applicability

This document outlines procedures for the extraction and subsequent determination of ammonium, sodium, and potassium ions in filter extracts. Analytical procedures outlined are specific to the Dionex Model DX-500 Ion Chromatograph.

1.2 Summary of Method

Cations in solution are separated when passed through a surface-sulfonated ion exchange resin due to the differing affinities of the cations for the active sites on the resin. After separation, the cations pass through a suppressor column which exchanges all anions for OH ions. Species are detected and quantified as their hydroxides by a conductivity meter. The eluent is sulfuric acid which yields deionized water when passed through the suppressor column.

In a laboratory evaluation of the accuracy of the method, spiked PM2.5 filter extracts and quality assurance/quality control samples were analyzed for sodium, ammonium, and potassium ions. The accuracy (expressed as %recovery) achieved using the subject method are presented in Table 1.

To test the precision of the method, PM2.5 filter extracts were analyzed in duplicate, and the blank extracting solution and a low-level QC sample were analyzed seven times each. The results are summarized in Table 2.

TABLE 1. ACCURACY VALUES FOR QA/QC SAMPLES AND SPIKED PM2.5 FILTER EXTRACTS

Analyte	QA/QC Sample Average % Recovery* (range)	Spiked Extract Average % Recovery* (range)
NH ₄ ⁺	99.0 (91.0 - 108.0) n = 59	99.4 (96.4 - 105.9) n = 14
Na ⁺	104.2 (100.0 - 107.7) n = 20	101.8 (97.0 - 105.1) n = 5
K ⁺	103.0 (96.6 - 106.3) n = 20	99.2 (97.0 - 102.7) n = 5

^{*%} Recovery = (concentration found/concentration expected) \times 100

Date: August 14, 2003

Page 4 of 10

TABLE 2. PRECISION VALUES FOR PM2.5 FILTER EXTRACTS, EXTRACTING SOLUTION, AND QC SAMPLES

Analyte	Sample type		
	PM _{2.5} Filter Extracts Average RSD** (range)	Blank Extracting Solution Average (Std Dev), n = 7	0.05 ppm QC Sample RSD, n = 7
NH ₄ ⁺	0.6 (0.1 - 1.6) for 13 duplicates	0.000 (0.000)	2.0
Na ⁺	8.0 (0.6 - 19.8) for 17 duplicates	0.001 (0.003)	13.3
K ⁺	1.1 (0.1 - 3.1) for 17 duplicates	0.000 (0.000)	2.0

^{**} RSD = relative standard deviation (the standard deviation divided by the average value and expressed as a percentage)

1.3 Health and Safety Warnings

The PM2.5 ion analysis operations do not involve unusual risks from electrical equipment or chemical exposures. Standard RTI laboratory health and safety precautions will be followed.

1.4 Cautions

Laboratory personnel should always wear clean clothes and wash hands thoroughly before performing filter handling and analysis procedures. The use of gloves is required and will minimize the potential for laboratory contamination.

1.5 Interferences

Large amounts of cations eluting close to the ions of interest will result in an interference. No interferences have been observed in extracts analyzed by RTI to date. If interferences are observed, several steps to increase separation can be taken, such as reducing eluent strength and/or flow rate.

1.6 Personnel Qualifications

Personnel employed to perform ion analysis operations will have at least a Bachelor's degree in a laboratory science, and will be trained by a supervisor before being allowed to process client samples for the $PM_{2.5}$ program.

1.7 Apparatus and Materials

- Tubes, centrifuge (Nalgene)
- Pipets an assortment of sizes
- Ultrasonic bath fitted with epoxy-coated test tube rack to hold centrifuge tubes
- Mechanical shaker
- Ion chromatograph (Dionex Model DX-500 with LC20 chromatography module, an IP20 or IP25 isocratic pump, a CD20 conductivity detector, a Dionex AS40 automated sampler and PeakNet Control Windows 98 Workstation) with Dionex CG12A cation guard column, Dionex CS12A cation separator column, and a Dionex CSRS-ULTRA cation self-regenerating suppressor
- Pressurized eluent reservoirs
- Volumetric flasks an assortment of sizes
- Dionex Autosampler Vials with filter caps

1.8 Reagents

Use ACS reagent grade chemicals and $18.2M\Omega$ -cm deionized water for the preparation of all solutions. Dry chemicals used for the preparation of calibration standards at 105° C for 2 hours and cool in a desiccator immediately before use.

- 1. Eluent, 22mN Sulfuric Acid: Dilute 22 mL 1N H₂SO₄ to 1 liter using deionized water. Sonicate for 15 minutes just prior to use to de-gas the solution.
- 2. Calibration Standard Stock Solution, 1000 mg/L each NH₄⁺, Na⁺, and K⁺: Dissolve 2.9654 g NH₄Cl, 2.5420 g NaCl, and 2.2284 g K₂SO₄ in 1 liter deionized water.

- 3. Standard Solution A: Dilute 10 mL stock solution to 100 mL with deionized water (100 mg/L NH₄⁺, Na⁺, K⁺).
- 4. Standard Solution B: Dilute 10 mL Standard Solution A to 100 mL with deionized water (10 mg/L NH₄⁺, Na⁺, K⁺).
- 5. Ethanol, nanopure reagent

1.9 Calibration

Using Standard Solutions A and B, prepare calibration standards with deionized water in 100 mL volumetric flasks as shown in Table 3. Prepare fresh calibration standards weekly.

TABLE 3. PREPARATION OF AMMONIUM ION CALIBRATION STANDARDS

Standard	NH ₄ ⁺ , Na ⁺ , K ⁺ (mg/L each)	mL of Standard Solution/100 mL		
STANDARD SOLUTION A				
1	25.0	25.0		
2	10.0	10.0		
3	3.0	3.0		
STANDARD SOLUTION B				
4	1.0	10.0		
5	0.3	3.0		
1 mg/L STANDARD (Standard 4)				
6	0.1	10.0		
7	0.05	5.0		

1.10 Sample Collection

Sample collection is not applicable to this SOP because samples are acquired by the state agency responsible for exposing the filters.

1.11 Sample Handling

Note: Additional information on this topic can be found in the SOP "Sample Receiving, Shipping, and Archiving Procedures for the PM_{2.5} Chemical Speciation Program," Research Triangle Institute, Center for Environmental Measurements and Quality Assurance, 1999.

RTI will provide chain-of-custody documentation with all sample shipments to track and ensure that samples are collected, transferred, stored, and analyzed by authorized personnel; sample integrity is maintained during all phases of sample handling and analysis; and an accurate written record is maintained of sample handling and treatment from the time of its collection, through the laboratory analytical process, to the eventual relinquishing of all data to the client.

Upon initial receipt of filters, RTI will prepare a Filter Inventory Sheet containing the filter identification numbers, box numbers, date received, date inspected and number of filters rejected. This form will allow laboratory personnel to select and use the filter boxes in the proper sequence.

1.12 Filter Extraction Procedure

1.12.1 Nylon Filters

NOTE: Filters to be analyzed for anions <u>and</u> cations or for cations only will be extracted with deionized water.

To extract the filters, the analyst will do the following:

- (1) Remove filters to be extracted from the freezer and allow them to equilibrate to room temperature.
- (2) Using gloved hands and tweezers, place each filter in a Nalgene centrifuge tube that has been labeled with the sample ID. (The label is carefully taped near the top of the Centrifuge tube to prevent loss during sonication.)
- (3) Label a 50-ml extraction tube as "Reagent Blank DI H₂O".
- (4) Add 25.0 mL of deionized water to each tube using a calibrated automatic pipette.
- (5) Ensure that the filter is completely submerged in the extraction solution.
- (6) Place the batch of centrifuge tubes in an epoxy-coated wire test tube rack and place the extraction rack in the ultrasonic bath. Make sure that the water level of the ultrasonic bath

PM2.5 Cation Analysis Revision 5 Date: August 14, 2003 Page 8 of 10

is higher than the extraction solution level but below the screw cap. Sonicate for 60 minutes.

CAUTION:	Monitor the bath temperature during sonication. The temperature
	should not exceed 27°C. Add ice initially to lower the temperature and
	add ice as necessary during the sonication to maintain an acceptable
	temperature.

- (7) Install the extraction racks on the mechanical shaker and shake overnight in a cold room (\leq 4°C) at approximately 60 cycles per minute.
- (8) Record the date of extraction on the RTI Sample Log Form.
- (9) Store the extracts in a refrigerator until analysis.

1.12.2 Teflon Filters

- (1) Remove filters to be extracted from the freezer and allow them to equilibrate to room temperature.
- (2) Using gloved hands and tweezers, place each filter in a Nalgene centrifuge tube that has been labeled with the sample ID. (The label is carefully taped near the top of the centrifuge tube to prevent loss during sonication.)
- (3) Label a 50-ml extraction tube as "Reagent Blank DI H₂O".
- (4) Remove the caps from all 50-ml extraction tubes. To prevent contamination place the caps in an upside-down position.
- (5) Using an Epppendorf 100- μ l pipette, wet the entire surface of each Teflon filter with 100 μ l of nanopure ethanol. This is done by very slowly pipetting the ethanol on the center of the filter. Capillary action will distribute the ethanol over the entire surface. The "Reagent Blank DI H₂O" tube will not contain a filter. Add the 100 μ l of ethanol directly to the bottom of the tube.

NOTE: Before proceeding, visually inspect each filter to be sure that the entire filter surface is wet.

(6) Using a calibrated automatic pipette, add 25.0 ml of deionized water to each extraction tube. The deionized water must have a resistance of at least $18.2M\Omega$ -cm.

- (7) Recap all extraction tubes tightly to prevent leakage during the extraction procedure. Be sure that the exposed area of the filter is completely immersed in the extraction solution.
- (8) Place the extraction rack in the ultrasonic bath. Make sure that the water level of the ultrasonic bath is higher than the extraction solution level but below the screw cap. Sonicate for 60 minutes.

CAUTION:	Monitor the bath temperature during sonication. The temperature
	should not exceed 27°C. Add ice initially to lower the temperature and
	add ice as necessary during the sonication to maintain an acceptable
	temperature.

- (9) Install the extraction racks on the mechanical shaker and shake overnight in a cold room (\leq 4°C) at approximately 60 cycles per minute.
- (10) Record the date of extraction on the RTI Sample Log Form.
- (11) Store the extracted filters in the refrigerator prior to analysis.

1.13 IC Procedure

- 1. Fill the eluent reservoirs with eluent.
- 2. Start the eluent flow, activate the self-regenerating suppressor, and allow the baseline to stabilize.
- 3. Inject two eluent blanks to flush the system and to ensure that the system is operating properly.
- 4. Using the calibration schedule, perform the daily multipoint calibration over the appropriate range followed by control/quality assurance (QC/QA) samples listed below:
 - A QC sample with NH₄⁺, Na⁺, and K⁺ concentrations typical of the higher concentrations found in actual filter extracts.
 - A QC sample with NH₄⁺, Na⁺, and K⁺ concentrations typical of the lower concentrations found in actual filter extracts.
 - A commercially prepared, NIST-traceable quality assurance sample containing known concentrations of NH₄⁺, Na⁺, and K⁺.

If the observed value for any cation differs by more than 10 percent from the known value, identify and correct the problem before analyzing samples.

- 5. Load the filter extracts into the autosampler vials according to the schedule prepared for that day. The daily schedule includes duplicate samples, spiked samples and quality control/quality assurance samples.
- 6. Begin the analysis run, occasionally checking to ensure that the system is operating properly.
- 7. Examine the data at the end of the run. If the NH₄⁺, Na⁺, or K⁺ concentration of any extract exceeds the upper end of its calibration curve, dilute the extract appropriately and analyze that day or include with the samples to be analyzed the following day.

1.14 Calculations and Data Reduction

Peak areas are entered into the computer where calculations are performed using a quadratic fit to the calibration data. The quadratic fit yields the following:

$$y_i = ax_i^2 + bx_i + c$$

where:

y = the calculated sulfate concentration, mg/L

x =the instrument response

The calibration curve from 0.05 to 10.0 ppm is used for the calculation of the extract NH_4^{+} , Na^+ , and K^+ concentrations. If a cation concentration exceeds 10 ppm, the extract is diluted appropriately (usually 5-fold) to bring the cation concentration into the calibration range and reanalyzed.

2.0 Quality Control and Quality Assurance

Compare the regression parameters (a, b, c and correlation coefficient) for the standard curves with those obtained in the past. If they exceed the control limits, stop the analysis and identify the problem.

Analyze quality control samples (see Section 1.13) at the beginning of every analytical run. Compare the results with those obtained during previous QC tests. If the observed concentration of any ion differs from the known value by greater than 10%, stop the analysis until the problem is identified and corrected. Analyze a duplicate sample, a QA/QC sample, and a spiked sample after at least every 20 field samples.

3.0 Reference

DRI Document No. 8068.1F4, Appendix D, Section 4.2